



Kentucky Department for Medicaid Services Drug Review and Options for Consideration

The following table lists the Agenda items scheduled, as well as the Options for Consideration, to be presented and reviewed at the **March 16, 2017** meeting of the Pharmacy and Therapeutics Advisory Committee.

Single Agent Reviews	Options for Consideration		
New Products to Market:	Non-prefer in the PDL class: Topical Antifungal Agents (Antifungals, Topical)		
DermacinRx® Therazole PakTM	 Non-prefer in the PDL class: Topical Antifungal Agents (Antifungals, Topical) Length of Authorization: 1 month DermacinRx® Therazole Pak™ (clotrimazole/betamethasone dipropionate packaged with zinc oxide) is a cream formulation of an azole-antifungal and a corticosteroid indicated for the topical treatment of symptomatic inflammatory tinea pedis, tinea cruris, and tinea corporis due to Epidermophyton floccosum, Trichophyton mentagrophytes, and Trichophyton rubrum in those ≥ 17 years of age. Available as a cream of 10 mg clotrimazole and 0.64 mg of betamethasone dipropionate. Criteria for Approval: Trial and failure of two different preferred agents; OR Is there any reason that the patient cannot be switched to a preferred medication? Document the details. Acceptable reasons include: Adverse reaction to preferred drugs Allergy to preferred drugs Contraindication to preferred drugs Age Limit = ≥ 17 years Quantity Limit = 180 grams per month (45 grams per week is the maximum 		
	usage per the package insert)		
New Products to Market: Vemlidy®	 Non-prefer in PDL class: Anti-infectives: Hepatitis B (Hepatitis B Agents) Length of Authorization: 6 months initial; 1 year renewal Vemlidy® (tenofovir alafenamide fumarate [TAF]) is a nucleoside analog reverse transcriptase inhibitor indicated for the treatment of chronic hepatitis B virus infection in adults with compensated liver disease. Available as a 25 mg tablet. Criteria for Approval: Diagnosis of Hepatitis B virus infection; AND Child-Pugh score is not B or C (decompensated cirrhosis); AND Not concurrently using any P-gp inducers (oxcarbazepine, phenobarbital, 		
	Child-Pugh score is not B or C (decompensated cirrho		



Single Agent Reviews	Options for Consideration				
	Not concurrently taking tenofovir disoproxil (Viread); AND				
	• Not HIV-1 positive using TAF as monotherapy.				
	Age Limit = ≥ 18 years Quantity Limit = 30 tablets per 30 days OR, if the patient is on				
	carbamazepine, then 60 tablets per 30 days.				
	*Note: Prior Authorization review and appropriate dosage to be determined by the Contact Center.				
New Products to Market:	Non-prefer in the PDL class: Oral Oncology, Other (Oncology Oral, Other)				
Rubraca TM	Length of Authorization: 6 months; may be renewed				
	 Rubraca™ (rucaparib) is a poly ADP-ribose polymerase (PARP) inhibitor indicated for use as single-agent therapy for treatment of adult females with advanced ovarian cancer that is associated with deleterious BRCA mutations in which patients have failed 2 or more other chemotherapies. Available as 200 mg and 300 mg tablets. 				
	Criteria for Approval:				
	Must have advanced disease; AND				
	Have a deleterious BRCA mutation as detected by an FDA-approved test (e.g., FoundationFocus CDxBRCA); AND				
	Must be used as a single agent; AND				
	• Must have received treatment with at least 2 prior lines of chemotherapy.				
	Age Limit = ≥ 18 years				
	Quantity Limit = 60 tablets per 30 days (1,200 mg per day is max dose)				
New Products to Market: BromSite TM	Non-prefer in the PDL class: Ophthalmic NSAIDs (Ophthalmics, Anti-inflammatories)				
	Length of Authorization: 3 weeks				
	• BromSite TM (bromfenac 0.075%) is a nonsteroidal anti-inflammatory				
	(NSAID) indicated for the treatment of postoperative inflammation and				
	prevention of ocular pain in patients undergoing cataract surgery.				
	Available as a 0.075% ophthalmic solution.				
	Criteria for Approval:				
	• Cataract surgery; AND				
	• Trial and failure of 1 preferred ophthalmic NSAID; OR				
	• Is there any reason that the patient cannot be switched to a preferred medication? Document the details. Acceptable reasons include:				
	 Adverse reaction to preferred drugs; 				
	- Allergy to preferred drugs; AND				
	 Contraindication to preferred drugs. 				
	Age Limit = ≥ 18 years				
New Products to Market:	Non-prefer in the PDL class: Platelet Aggregation Inhibitors				
Yosprala TM	Length of Authorization: 1 year				



Single Agent Reviews	Options for Consideration		
	• Yosprala TM (aspirin/omeprazole) is a combination of aspirin (an antiplatelet) and omeprazole (a Proton Pump Inhibitor [PPI]) indicated for patients who require aspirin for secondary prevention of cardiovascular and cerebrovascular events who are at risk of developing aspirinassociated gastric ulcers. It is not interchangeable with the individual components of aspirin and omeprazole. Available as 325 mg delayedrelease aspirin/40 mg immediate-release omeprazole or as 81 mg delayedrelease aspirin/40 mg immediate-release omeprazole.		
	Criteria for Approval:		
	• Has the patient had a therapeutic trial and treatment failure of at least 1 preferred drug? Document the details; OR		
	• Is there any reason that the patient cannot be switched to a preferred medication? Document the details. Acceptable reasons include:		
	 Adverse reaction to preferred drugs; 		
	 Allergy to preferred drugs; AND 		
	 Contraindication to preferred drugs. Limitations of Use: Not for use as initial dose of aspirin therapy during onset 		
	of acute coronary syndrome, acute myocardial infarction, or before		
	percutaneous coronary intervention. It has not been shown to reduce the risk of gastrointestinal bleeding due to aspirin. Quantity Limit: 1 tablet per day		



Full Class Reviews	Options for Consideration		
	Agents in the following Therapeutic Classes are subject to status changes from what is on the current Preferred Drug List (PDL).		
Anticoagulants	DMS to select preferred agent(s) based on economic evaluation; however, at least 1 low molecular weight heparin, 1 factor Xa inhibitor, and 2 oral anticoagulants should be preferred.		
	• Agents not selected as preferred will be considered non-preferred and require PA.		
	• For any new chemical entity in the <i>Anticoagulants</i> class, require PA until reviewed by the P&T Advisory Committee.		
Antifungals, Oral	DMS to select preferred agent(s) based on economic evaluation; however, at least fluconazole, griseofulvin, nystatin, and terbinafine should be preferred.		
	Agents not selected as preferred will be considered non-preferred and require PA.		
	• For any new chemical entity in the <i>Antifungals, Oral</i> class, require PA until reviewed by the P&T Advisory Committee.		
Cephalosporins & Related	1st Generation:		
(Antibiotics: Cephalosporins 1st, 2nd, 3rd Generation)	DMS to select preferred agent(s) based on economic evaluation; however, at least cephalexin should be preferred.		
	Agents not selected as preferred will be considered non-preferred and require PA.		
	• For any new chemical entity in the <i>1st Generation Cephalosporin</i> class, require PA until reviewed by the P&T Advisory Committee.		
	2 nd Generation:		
	DMS to select preferred agent(s) based on economic evaluation; however, at least cefuroxime should be preferred.		
	Agents not selected as preferred will be considered non-preferred and require PA.		
	• For any new chemical entity in the <i>2nd Generation Cephalosporin</i> class, require PA until reviewed by the P&T Advisory Committee.		
	3 rd Generation:		
	DMS to select preferred agent(s) based on economic evaluation; however, at least cefixime and cefpodoxime should be preferred.		
	Agents not selected as preferred will be considered non-preferred and require PA.		
	• For any new chemical entity in the 3rd Generation Cephalosporin class, require PA until reviewed by the P&T Advisory Committee.		



Full Class Reviews	Options for Consideration			
GI Motility, Chronic	DMS to select preferred agent(s) based on economic evaluation; however,			
(GI Motility Agents)	at least 1 unique chemical entity should be preferred.			
	• Agents not selected as preferred will be considered non-preferred and will require PA.			
	• For any new chemical entity in the <i>GI Motility, Chronic</i> class, require PA until reviewed by the P&T Committee.			
Hypoglycemics, Incretin	Amylin Analogs:			
Mimetics & Enhancers	DMS to select preferred agent(s) based on economic evaluation.			
(Diabetes: Amylin Analogs,	Allow for use of pramlintide with active insulin therapy only.			
DPP-4 Inhibitors, GLP-1 Receptor Agonists)	Agents not selected as preferred will be considered non-preferred and will require PA.			
	• For any new chemical entity in the <i>Hypoglycemics, Amylin Analogues</i> class, require PA until reviewed by the P&T Advisory Committee.			
	DPP-4 Inhibitors:			
	DMS to select preferred agent(s) based on economic evaluation; however, at least 1 single entity agent should be preferred.			
	• Agents not selected as preferred will be considered non-preferred and will require PA.			
	• For any new chemical entity in the <i>Hypoglycemics, DPP4-Inhibitors</i> class, require PA until reviewed by the P&T Advisory Committee.			
	GLP-1 Receptor Agonists:			
	New addition to the class: <u>AdlyxinTM</u>			
	Non-prefer in this class. Length of Authorization: 1 year			
	• Adlyxin TM (lixisenatide) is a glucagon-like peptide-1 (GLP-1) receptor agonist administered subcutaneously once daily within 1 hour of the first meal of the day, indicated as an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus. Available as 50 mcg/ mL and 100 mcg/ mL solution in a 3 mL prefilled pen.			
	Criteria for Approval:			
	Diagnosis of type 2 diabetes mellitus; AND			
	Trial and failure of metformin, document; AND			
	Trial and failure of a preferred GLP-1 receptor agonist.			
	Age Limit = ≥ 18 years			
	Quantity Limit = 2 pens per 28 days			
	New addition to the class: Soliqua™			
	Non-prefer in this class.			
	Length of Authorization: 1 year			
	• Soliqua TM (insulin glargine/lixisenatide) is a fixed-dose combination of insulin glargine (Lantus [®]) and the GLP-1 agonist, lixisenatide (Adlyxin TM)			



Full Class Reviews	Options for Consideration			
	administered subcutaneously once daily within 1 hour of the first meal of the day, indicated as an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus who are not controlled with basal insulin (< 60 units) or lixisenatide. Available as 100 unit insulin glargine/ 33 mcg lixisenatide per mL solution in a 3 mL prefilled multidose pen.			
	Criteria for Approval:			
	Diagnosis of type 2 diabetes mellitus; AND			
	Trial and failure of lixisenatide or basal insulin separately; AND			
	Trial and failure of preferred GLP-1 receptor agonists and preferred long- acting insulin; AND			
	Not used in combination with other GLP-1 agonists.			
	Age Limit = ≥ 18 years			
	Quantity Limit = 5 pens (1 carton) per 25 days			
	New addition to the class: Xultophy®			
	Non-prefer in this class.			
	Length of Authorization: 1 year			
	• Xultophy® (insulin degludec/liraglutide) is a fixed-dose combination of insulin degludec (Tresiba®) and the GLP-1 agonist, liraglutide (Victoza®) administered subcutaneously once daily at the same time of day, with or without food, indicated as an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus who are not controlled on basal insulin (< 50 units daily) or liraglutide (≤ to 1.8 mg daily).			
	Criteria for Approval:			
	Diagnosis of type 2 diabetes mellitus; AND			
	Trial and failure of liraglutide or basal insulin; AND			
	Trial and failure of preferred GLP-1 receptor agonists and insulin; AND			
	• Not used in combination with other GLP-1 agonists.			
	Age Limit = ≥ 18 years			
	Quantity Limit = 5 pens (1 carton) per 30 days			
	DMS to select preferred agent(s) based on economic evaluation; however, at least one unique chemical entity should be preferred.			
	• Continue to require PA for all agents in this class to ensure appropriate utilization.			
	• For any new chemical entity in the <i>Hypoglycemics, GLP-1 Receptor Agonists</i> class, require PA until reviewed by the P&T Advisory Committee.			



Full Class Reviews	Options for Consideration					
Hypoglycemics, Insulins & Related	DMS to select preferred agent(s) based upon economic evaluation; however, at least 1 insulin per class should be preferred.					
(Diabetes: Injectable Insulins)	 Agents not selected as preferred will be considered non-preferred and will require PA. For any new chemical entity in the <i>Hypoglycemics, Insulins and Related</i> 					
	class, require PA until reviewed by the P&T Advisory Committee.					
Hypoglycemics, SGLT2s	New addition to the class: Invokamet® XR					
(Diabetes: SGLT2	Non-prefer in this class.					
Inhibitors)	Length of Authorization: 6 months initial; 1 year renewal					
	• Invokamet® XR (canagliflozin/metformin) is a sodium-glucose cotransporter 2 (SGLT2) inhibitor and biguanide combination product indicated as an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus when treatment with both canagliflozin and metformin is appropriate. Available as 50 mg/ 500 mg, 50 mg/ 1000 mg, 150 mg/ 500 mg, and 150 mg/ 1000 mg extended-release tablets.					
	Criteria for Approval:					
	Diagnosis of type 2 diabetes mellitus; AND					
	• Documented reason Invokamet® cannot be used (Invokamet® is preferred without PA).					
	Quantity Limit = 2 extended-release tablets per day.					
	• DMS to select preferred agent(s) based on economic evaluation; however, at least 1 unique chemical entity should be preferred.					
	• Agents not selected as preferred will be considered non-preferred and will require PA.					
	• For any new chemical entity in the <i>Hypoglycemics, SGLT2 Inhibitors</i> class, require PA until reviewed by the P&T Advisory Committee.					
Hypoglycemics, Sulfonylureas	• DMS to select preferred agent(s) based on economic evaluation; however, at least 2 unique second generation sulfonylureas should be preferred.					
(Diabetes: Sulfonylureas)	• Agents not selected as preferred will be considered non-preferred and will require PA.					
	• For any new chemical entity in the <i>Hypoglycemics, Sulfonylureas</i> class, require PA until reviewed by the P&T Advisory Committee.					
Tetracyclines (Antibiotics: Tetracyclines)	DMS to select preferred agent(s) based on economic evaluation; however, at least generic formulations of doxycycline and minocycline should be preferred.					
	• If demeclocycline is selected as non-preferred, allow for its use in SIADH only.					
	Agents not selected as preferred will be considered non-preferred and require PA.					



Full Class Reviews	Options for Consideration			
	• For any new chemical entity in the <i>Tetracyclines</i> class, require PA until reviewed by the P&T Advisory Committee.			
Orkambi® Criteria Review	v Current Criteria:			
	Length of Authorization: 6 months; may be renewed			
	Criteria for Approval:			
	• Diagnosis of cystic fibrosis homozygous for the F508del mutation in the CFTR gene confirmed by an FDA-approved CF mutation test; AND			
	● Baseline ophthalmic examinations if patient is 12 − 18 years of age.			
	Renewal Criteria;			
	• Stable or improved FEV ₁ ; AND			
	• Serum ALT or AST \leq 5 times the ULN, or ALT or AST, \leq 3 times the ULN with bilirubin \leq 2 times the ULN.			
	Age Limit = ≥ 12 years			
	Recommended Changes:			
	Age Limit = ≥ 6 years			
	Quantity Limit = 112 tablets per 28 days.			
	• Patient age 6 – 11 years = 2 tablets orally every 12 hours with fat-			
	containing food. Use the lumacaftor 100 mg/ivacaftor 125 mg tablet			
	strength.			
	• Patient age ≥ 12 years = 2 tablets orally every 12 hours with fat-containing			
	food. Use the lumacaftor 200 mg/ ivacaftor 125 mg tablet strength. Renewal Criteria:			
	Patient has not received a lung transplant; AND			
	No unacceptable toxicity from the drug; AND			
	 Disease response as indicated by 1 or more of the following; 			
	 Disease response as indicated by 1 of more of the following; Decreased pulmonary exacerbations as compared to pretreatment 			
	baseline			
	Improvement or stabilization of lung function compared to baseline			
	Decrease in decline of lung function			
	 Improvement in quality of life, weight gain, or growth 			



Consent Agenda	Options for Consideration		
	For the following therapeutic classes, there are no recommended changes to the currently posted Preferred Drug List (PDL) status.		
	Antibiotics, GI	•	Hypoglycemics, Meglitinides
	• Antibiotics, Inhaled	•	Hypoglycemics, Metformins
	Antibiotics, Vaginal	•	Hypoglycemics, Thiazolidinediones
	Antipsoriatics, Topical	•	Ketolides/Macrolides
	COPD Agents	•	Oxazolidinones
	• Fluoroquinolones, Oral	•	Penicillins
	• Hypoglycemics, Alpha-Glucosidase Inhibitors	•	Sulfonamides, Folate Antagonists
	For the following therapeutic classes, there are no recommended changes other than a brand/generic switch.		
	N/A		

